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(54) Title: PRODUCTION OF ANTIBODIES USING CRE-MEDIATED SITE-SPECIFIC RECOMBINATION

(57) Abstract

A nethod to produce a cell expressing a mithody from a general exquence of a cell which has a modified immunopidabile locus is disclosed. The restend sizes advanced of Co-medizated shee opends: recombination and floored-tops recombination. The method involves fars immoducing a for size into an immunopidabilin locus and then using the lot size as a target for immoducion of gene modifications via Cre-medizated in vive recombination.

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PRODUCTION OF ANTIBODIES USING CRE-MEDIATED SITE-SPECIFIC RECOMBINATION

Technical Field

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The invention relates to a process to produce antibodies from genetic loci modified using recombinant DNA vectors and site-specific recombination leading to the production of modified antibody molecules by transfected cells. More particularly, the invention relates to the use of Cre-mediated site-specific recombination for modifying immunoglobulin loci, for instance, to replace all or a portion of either the constant region or variable region of an antibody molecule to form a modified antibody molecule. One particular aspect relates to class-switching of antibody genes in antibody-producing lymphoid cells in situ whereby a constant region of an immunoglobulin gene is replaced with a constant region of another class, thereby producing a modified antibody with a changed isotype. Another aspect relates to modification of the variable region, or a portion thereof, which is replaced or exchanged with a variable region having a different or altered antigen specificity.

Background Art

The basic immunoplobulin structural unit in vertebrate systems is composed of two identical "light" polypeptide chains of molecular weight approximately 23,000 daitons, and two identical "heavy" chains of molecular weight 53,000-70,000. The four chains are joined by disulfide bonds in a "" configuration in which the light chains bracket the heavy chains starting 5

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at the mouth of the Y and continuing through the divergent region or "branch" portion which is designated the Fab region. Heavy chains are classified as gamma (y), mu (μ) , alpha (α) , delta (δ) , or epsilon (ϵ) , with some subclasses among them which vary according to species; and the nature of this chain, which has a long constant region, determines the "class" of the antibody as IgG, IgM, IgA, IgD, or IgE, respectively. Light chains are classified as either kappa (K) or lambda (λ) . Each heavy chain class can be associated with either a kappa or lambda light chain. The light and heavy chains are covalently bonded to each other, and the "tail" portions of the two heavy chains are bonded to each other by covalent disulfide linkages when the immunoglobulins are generated either by hybridomas or by B calls.

The amino acid sequence of each immunoglobulin chain runs from the N-terminal end at the top of the Y to the C-terminal end at the bottom. The N-terminal end contains a variable region (V) which is specific for the antigen to which it binds and is approximately 100 amino acids in length, there being variations between light and heavy chain and from antibody to antibody. The variable region is linked in each chain to a constant region (C) which extends the remaining length of the chain. Linkage is seen, at the genomic level. as occurring through a linking sequence known as the joining (J) region in the light chain gene, which encodes about 12 amino acids, and as a combination of diversity (D) region and joining (J) region in the heavy chain gene, which together encode approximately 25 amino acids. The remaining portions of the chain, the constant regions, do not vary within a particular class with the specificity of the antibody (i.e., the antiqen to which it binds). The constant region or class

determines subsequent effector function of the antibody, including activation of complement and other cellular responses, while the variable region determines the antigen with which it will react.

Since the development of the cell fusion technique for the production of noncional antibodies by Kohler and Milstein, many individual immunoglobulin species have been produced in quantity. Most of these sonoclonal antibodies are produced in a murine system and, therefore, have limited utility as human therapeutic agents unless modified in some way so that the murine monoclonal antibodies are not "recognized" as foreign epitopes and "neutralized" by the human immune system.

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One approach to this problem has been to attempt to develop human or "humanized" monoclonal antibodies, which are "recognized" less well as foreign spitopes and may overcome the problems associated with the use of monoclonal antibodies in humans. Applications of human s cell hybridoms-produced monoclonal antibodies hold great promise for the treatment of cancer, viral and microbial infections, 3 cell immunodeficiencies with diminished antibody production, and other diseases and disorders of the immune system.

However, several obstacles exist with respect to the development of human monoclonal antibodies. For example, with respect to monoclonal antibodies which recognise human tumor antigens for the diagnosis and treatment of cancer, many of these tumor antigens are not recognized as foreign antigens by the human immune system and, therefore, these antigens may not be immunogenic in man.

Another problem with human monoclonal antibodies is that most such antibodies obtained in cell culture are of one class or isotype, the IgM type. Under certain

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circumstances, monoclonal antibodies of one isotype might be more preferable than those of another in terms of their diagnostic or therapeutic efficacy since, as noted above, the isotype determines subsequent effector function of the antibody, including activation of complement and other cellular responses. For example, from studies on antibody-mediated cytolysis it is known that unmodified mouse monoclonal antibodies of subtype y2a and y3 are generally more effective in lysing target cells than are antibodies of the Yl isotype. This differential efficacy is thought to be due to the ability of the y2a and y3 subtypes to more actively participate in the cytolytic destruction of the target calls. Particular isotypes of a murine monoclonal antibody can be prepared either directly, by selecting from the initial fusion, or secondarily, from a parental hybridoma secreting monoclonal antibody of a different isotype, by using the "sib selection" technique to isolate class-switch variants (Steplewski et al., 1985, Proc. Natl. Acad. Sci. USA 82:8653; Spira et al., 1984, J Immunological Methods 74:307.

when human monocional antibodies of the 1g6 type are desired, however, it has been necessary to use such tedious techniques as cell sorting, to identify and isolate the few cells which are producing antibodies of the 1g6 or other type from the majority producing antibodies of the 1g8 type. A need therefore exists for an efficient method of switching antibody classes in isolated antibody-producing cells for any given antibody of a predeteximate of desired antipenic specificity.

Various solutions to these problems with monoclonal annual models for human use have been developed based on recent methods for the introduction of DNA into mammalian cells to obtain expression of immunoglobulin censes, particularly for production of othersic

inmunoglobulin molecules comprising a human and a non-human portion. More specifically, the antigen combining (wariable) region of the chimeric antibody is derived from a non-human source (e.g., murine), and the constant region of the chimeric antibody (which confers biological effector function to the immunoglobulin) is derived from a human source. Such "humannized" chimeric antibodies should have the antigen binding specificity of the non-human antibody molecule and the effector function conferred by the human antibody molecule.

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Generally, chimeric antibodies have been produced conventionally by procedures comprising the following steps (although not necessarily in this order): (1) · identifying and cloning the gene segment encoding the antigen binding portion of the antibody molecule: this gene segment (VDJ for heavy chains or VJ for light chains, or more simply, the variable region) may be obtained from either a cDNA or genomic source; (2) cloning the gene segments encoding the constant region or desired part thereof; (3) ligating the variable region with the constant region so that the complete chimeric antibody is encoded in a transcribable and translatable form; (4) ligating this construct into a vector containing a selectable marker and appropriate gene control regions; (5) amplifying this construct in bacteria: (6) introducing the DNA into eukaryotic cells (by transfection), most often cultured mammalian cells such as lymphocytes; (7) selecting for cells expressing the selectable marker; (8) screening for cells expressing the desired chimeric antibody; and (9) testing the antibody for appropriate binding specificity and effector functions.

Antibodies of several distinct antigen binding specificities have been manipulated by these procedures to produce chimeric proteins. In addition several

different effector functions have been achieved by linking new sequences to those snooding the antigen binding region. Some of these include enzymes (Neuberger et al., 1984, Meture 121:604), immunoglobulin constant regions from another species and constant regions of another immunoglobulin chain (Sharon et al., 1984, Mature 102:364; Tan et al., 1985, J. Immunol. 113:3563-3567). Neuberger et al., PCT Publication Wo 86/0153) (1984) also discloses production of chimeric antibodies and mentions, among the technique's many uses, the concept of "Class switching."

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Cabilly, et al., U.S. Patent No. 4,816,567 issued March 28, 1889, describes altered and native immunoglobulins, including constant-variable region chimeras, which are prepared in recombinant cell culture. The immunoglobulins contain variable regions which are immunologically capable of binding predetermined antigens. The vactors and methods disclosed are suitable for use in various host cells including a vide range of prokaryotic and sukaryotic organisms.

Fell et al., U.S. Patent No. 5,202,238 issued April 19,193, describes a process for producing chineric antibodies using recombinant DNA vectors and homologous recombination in vivo. The process uses novel recombination NA vectors to enginer targeted gene modification accomplished via homologous recombination in either (a) cell lines that produce antibodies having desired antigem specificities, so that the antigen combining site of an antibody molecule remains unchanged, but the constant region of the antibody molecule, or a portion thereof, is replaced or modified; or (b) cell lines that produce antibodies of desired classes which may demonstrate desired effector functions, so that the constant region of an antibody process so that the constant region of an antibody

molecule remains unchanged, but the variable region of the antibody molecule or a portion thereof, is replaced or modified. The reported efficiency of recombination was relatively low, however, ranging from 0.39% to 0.75% in several attempts to replace a mouse heavy chain constant region with a human counterpart, even when a selectable marker gene was used to recover recombinant commons.

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function.

Kucherlapati et al., in PCT Publication W091/10741 (published July 25, 1991) and in PCT Publication WO 94/02602 (published February 3, 1994) disclose xenogeneic specific binding proteins or antibodies produced in a non-primate viable mammalian host by immunization of the mammalian host with an appropriate immunogen. In particular these publications disclose production of antigen-specific human monoclonal antibodies from mice engineered with loci for human immunoglobulin heavy and light chains using yeast artificial chromosomes (YACs). Such mice produce completely human antibodies in response to immunization with any antigen normally recognized by the nouse immune system, including human antigens; and B-cells from these mice are used to make hybridomas producing human monoclonal antibodies via conventional hybridoma production methods. While production of completely human antibodies from transgenic murine hybridomas solves many of the previous problems of producing human monoclonal antibodies, it may be desirable to modify the loci in the cells which produce the antibodies, for instance, to enhance expression or to alter an effector

Sauer et al, U.S. Patent No. 4,959,317, issued September 25, 1990, describes a method for producing site-specific recombination of DNA in eukaryotic cells at sequences designated low sites. DNA sequences

comprising first and second low sites are introduced into aukaryotic cells and interacted with a recombinase designated Cre (typically, by transient expression of the recombinase from a plasmid), thereby producing recombination at the low sites. Exemplified eukaryotic cells included yeast cells and smoolayer cultures of a mouse cell line. Frequencies of Cre-nediated recombination ranged from 2-3t to 22% for repeated recombination tanged from 2-3t to 22% for repeated recombination attempts between an exemplary virus and a plasmid, and 98% for the case of deletion of a yeast leuz gene flamked by low sites. Bowever, neither recombination in lymphoid cells nor manipulation of immunoglobulin loci is disclosed by Sauer et al. Qu et al., 1993, Cell 21:1155-114 describes a

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method to generate a mouse strain in which the J region and the intron enhancer in the heavy-chain locus are deleted from embryonic stem cells using Cre-mediated size-specific recombination. They then analyzed the immunoglobulin isotypes formed by recombination in heterorygous mutant B cells, activated with LPS plus ILA. The authors used Cre-mediated size-specific recombination marely to modify the heavy chain locus in stem cells which are not antibody-producing cells, and only to modify that locus by deletion, rather than to produce new combinations of gene sequences in antibody-producing cells to produce chimeric or modified anythodies.

Johnson et al., in PCT Publication WO93/19172 (published September 9, 1993), and Waterhouse et al., in the journal article, Winciel Arids Res (1993) 21:2265-2266, describe use of Cre-mediated site-specific recombination to effect recombinations for creation of a combinatorial library of antibodies in phage vectors. In the latter publication, Johnson, Waterhouse and coworkers describe two explary phage vectors

designated A and B, where A encodes the light chain of a first antibody (and the heavy chain from a different antibody) and B encodes the heavy chain of the first antibody. In both vectors the variable heavy chain (VH) genes are flanked by two loxP sites, one of which is a mutant lowP site which prevents recombination within the vector from merely excising the VH genes. When Cre recombinase is provided in vivo by infecting the E. coli with a phage expressing Cre, vectors A and B can cointegrate by recombination between either mutant or wild-type loxP sites to create chimeric plasmids. Further recombination can then occur between the two wild-type or the two mutant loxP sites, to generate original vectors A and B or two new vectors, E and F. The heavy chains of A and B are therefore exchanged in E and F, and E encodes the Fab fragment of the first antibody for display as a fusion to the N-terminus of the phage gene 3 protein (g3p). The authors indicate that the method should allow the creation of extremely large combinatorial repertoires of phage-expressing antibodies, for example by providing a light chain repertoire in phage vector A and a heavy chain repertoire in phage vector B. However, these two publications do not disclose use of Cre-mediated sitespecific recombination for any other purpose besides exchange of heavy chain genes in the particular bacteriophage vectors disclosed and, more specifically, do not propose any modification of immunoglobulin sequences in the genome of any antibody-producing cell. Accordingly, the present invention provides for a novel use of Cre-mediated site-specific recombination for modification of immunoglobulin loci in the cellular

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genome of any antibody-producing cell.

Disclosure of the Invention

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The invention is directed to processes for producing a cell expressing a desired antihody from a genomic sequence by modification of an immunoglobulin locus using Cre-mediated site-specific recombination. The methods provide, inter alia, the advantage of increased efficiency of recombination over previous methods of modifying immunoglobulin loci directly in antibody-producing cells using only homologous recombination for modifying the loci.

Thus, in one aspect, the invention is directed to a method to produce a cell expressing an antibody from a gamonic sequence of the cell comprising a modified immunoplobulin locus. The invention method uses Cremediated site-specific recombination and comprises:

- (a) transfecting an antibody-producing cell with a first homology-targeting vector comprising: (1) a first lox site, and (ii) a targeting sequence homologous to a first DNA sequence adjacent to the region of the immunoglobulin locus of the genomic sequence which is to be converted to a modified region, so that the first lox site is inserted into the genomic sequence via sitespecific homologous recombination with genomic DNA in vivo:
- (b) transfecting said cell with a low-targeting vector comprising: (i) a second low site suitable for Cre-mediated recombination with the first low site, and (ii) a modifying sequence to convert said region of the immunol/bulin locus to a modified region;
- (c) interacting the low sites with Cre, so that the modifying sequence inserts into the genomic sequence via Cre-mediated site-specific recombination of the lox sites, thereby converting the region of the impunoplobulin locus to the modified region; and

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(d) selecting a transfectant in which the region of the immunoglobulin locus is converted to the modified region and which produces the antibody molecule.

In one preferred embodiment of this method, the lor-targeting vector further comprises a selectable marker gene operably linked to control regions such that the marker gene is expressed in the cell. On interacting the low sites with Cre, the marker gene inserts into the genomic sequence with the modifying sequence via Cre-mediated site-specific recombination of the low sites. In this embodiment, selecting for a transfectant comprises selecting for expression of the marker cene.

A second preferred embodiment of the method of the invention further comprises an additional step before step (b), which additional step comprises transfecting the cell with a second homology-targeting vector comprising: (i) a third lox site suitable for Cremediated recombination with the first and second lox sites, (ii) a first selectable marker gene operably linked to control regions such that the first marker gene is expressed in the cell, and (iii) a targeting sequence homologous to a second DNA sequence adjacent to the region of the immunoglobulin locus of the genomic sequence which is to be converted. In this embodiment the gene which is to be converted is flanked by the first and second DNA sequences, and the third lox site and first marker gene are inserted into the genomic sequence via site-specific homologous recombination with genomic DNA in vivo.

Also in this second preferred embodiment the second targeting vector further comprises a second selectable marker gene operably linked to control regions such that the second marker gene is expressed in the cell. On interacting the low sites with Cre, the

second marker gene inserts into the genomic sequence with the modifying sequence via Cre-mediated site-specific recombination of appropriate lox sites. In this smbodiment selecting for a transfectant comprises selecting for expression of the second marker gene. In a further variation of this embodiment of the method of the invention, on interacting the lox sites with Cre, after the modifying sequence and second marker gene insert into the genomic sequence, the first marker gene and the region to be converted are delated by Cremediated site-specific recombination, and selecting for a transfectant further comprises selecting for a transfectant further comprises selecting for a transfectant further comprises.

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In some embodiments of the invention method, the modifying sequence comprises a regulatory nucleotide sequence which replaces all or a portion of the regulatory sequences of the immunoglobulin genes of the genomic sequence to provide modified expression of those immunoglobulin genes. In other embodiments, the modifying sequence comprises a nucleotide sequence that encodes a translation product to replace all or a portion of either the constant region or the variable region of an antibody molecule to form a modified antibody molecule. In some of the latter embodiments, the region of the immunoglobulin locus to be converted to a modified region comprises a constant region gene and the modifying sequence comprises a nucleotide sequence that encodes a translation product to replace all or a portion of the constant region of the antibody produced by said constant region gene with a modified constant region.

Sometimes the constant region gene of the genomic sequence is a human constant region gene and the modified constant region gene encodes a different human constant region. The modified constant region gene may PCT/US96/04445

comprise a light chain gene or a heavy chain gene. In various embodiments of the invention method to produce a cell expressing an antihody molecule with a modified constant region, the modified constant region gene comprises a sequence encoding an enzyma, toxin, hormone, growth factor, linker or mutant constant region with an attered heavy chain effector function.

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In some embodiments of the invention method, the cupressing an antibody molecule is a cell of a lymphoid cell line. This lymphoid cell line may be a murine hybridoma cell line producing either a murine, human or chimeric antibody. In some embodiments, the hybridoma cell line is producing a human antibody by expression of human immunoglobulin genes. In one particular embodiment the cell is a murine lymphoid cell producing a human antibody by expression of human

particular embodiment the call is a murne symptotic exproducing a human antibody by expression of human immunoglobulin gense. In one variation of this embodiment, the constant region gene of the genomic sequence is a human constant region (c) gene of the mu class, i.e., a c, gene, and the modifying sequence comprises a human c gamam (c), constant region gene.

In another aspect the invention is directed to a entry especial anniholdy from a genomic sequence of the cell comprising a modified immunoglobulin locus, where the cell is produced by the invention method above.

Another aspect of the invention is a method to produce a cell expressing an antibody nolecule from a genomic sequence of the cell with a lor site adjacent to or integrated within a region of the immunoglobulin locus encoding the antibody nolecule, for use in modifying that region using Cre-mediated site-specific recombination. This method comprises: (a) transfecting an antibody-producing cell with a first targeting vector comprising: (i) a first lor site, and (ii) a targeting

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invention.

sequence homologous to a first DNA sequence adjacent to the region of the immunoglobulin locus of the genomic sequence which is to be converted to a modified region, so that the first low site is inserted into the genomic sequence via site-specific homologous recombination with genomic DNA in vivo. This method further comprises (b) selecting a transfectant with the lox site inserted into the genomic DNA adjacent to the region which is to be converted. This selection may be achieved using one or more selectable marker genes in a vector used to insert a lox site into the genomic sequence via site-specific homologous recombination with genomic DNA in vivo. Another facet of this aspect of the invention relates to a cell expressing an antibody molecule with a lox site adjacent to or integrated within a region of the immunoglobulin locus encoding an antibody molecule. where the cell is produced by the above method of the

In yet another aspect the invention is directed to an embryonic stem cell of a non-primate marmal comprising a genome comprising a transgenic non-primate mammal comprising a genome comprising at least a functional portion of a human heavy chain immunoqiboulin locus or at least a functional portion of a human light chain immunoqiboulin locus. In the genome of this stem cell, a lox site is adjacent to or integrated within a region of said human heavy chain or light chain immunoqiboulin locus. In a praferred embodiment, this stem cell is a murine stem cell.

A related aspect of the invention is a transgenic non-primate manual comprising a genome comprising at least a functional portion of a human heavy chain immunoglobulin locus or at least a functional portion of a human light chain immunoglobulin locus. In the genome of this stem cell, a low site is adjacent to or

integrated within a region of said human heavy chain or light chain immunoglobulin locus. In a preferred embodiment, this transgenic mammal is murine.

Other aspects of the invention are described below.

Brief Description of the Drawings

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Figure 1 represents a schematic diagram of one embodiment (Scheme A) of the invention method to produce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated site-specific recombination. This scheme requires insertion of only a single low site adjacent to the constant region gene to be modified.

Figure 2 represents a schematic diagram of another embodisems (Scheme 8) of the invention method to produce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated site specific recombination. This scheme uses two low sites integrated into the genome of the antibody-producing cell, flanking the constant region to be modified.

cell, flanking the constant region to be modified. Figure) represents a schematic diagram of one embodiment (Scheme C) of the invention method to produce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated sitespecific recombination. This scheme also requires insertion of only a single low site adjacent to the constant region gene to be modified. However, the low site is inserted into the genome of a transpenic animal from which the cell expressing the antibody is obtained.

Modes of Carrying Out the Invention

Definitions

The following terms, as used herein, whether in the singular or plural, shall have the meanings indicated:

encodes all or a portion of either the constant region or variable region of an antibody molecule or all or a portion of a regulatory nucleotide sequence that controls expression of an antibody molecule. Immunoglobulin loci for heavy chains may include but are not limited to all or a portion of the V, D, J, and switch regions (including intervening sequences called introns) and flanking sequences associated with or adjacent to the particular heavy chain constant region gene expressed by the antibody-producing cell to be transfected and may include regions located within or downstream of the constant region (including introns). Immunoglobulin loci for light chains may include but are not limited to the V and J regions, their upstream flanking sequences, and intervening sequences (introns). associated with or adjacent to the light chain constant region gene expressed by the antibody-producing cell to be transfected and may include regions located within or downstream of the constant region (including introns). Immunoglobulin loci for heavy chain variable regions may include but are not limited to all or a portion of the V, D, and J regions (including introns) and flanking sequences associated with or adjacent to the particular variable region gene expressed by the antibody-producing cell to be transfected. Immunoglobulin loci for light chain variable regions may include but are not limited

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Immunoglobulin locus: a nucleotide sequence that

Modified Antibody: an antibody molecule in which (a) the constant region, or a portion thereof, is altered, replaced or exchanged so that the antigen binding site (variable region) is linked to a constant

to the V and J region (including introns) and flanking sequences associated with or adjacent to the light chain variable region gene expressed by the antibody-producing

cell to be transfected.

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region of a different or altered class, effector function and/or species, or an entirely different nolecule which confers new properties to the modified antibody, e.g., an enyme, toxin, hormone, growth factor, drug, and the like; or (b) the variable region, or a portion thereof, is altered, replaced or exchanged with a variable region having a different or altered antigen specificity.

Modifying Sequence: a nucleotide sequence that encodes (a) a translation product to modify or replace all or a portion of either the constant region or variable region of an antibody molecule to form a modified antibody molecule, or (b) one or more regulatory nucleotide sequences to modify or replace all or a portion of the regulatory sequences of an immunoglobulin locus sequence to provide modified expression of that immunoglobulin locus sequence. This regulatory sequence may include, for instance, a promoter, enhancer, an intron, a switch sequence, a ribosome binding site, and the like, and may be in addition to or instead of a sequence that encodes a translation product to replace a region of an antibody molecule. Suitable regulatory nucleotide sequences are known in the art. The regulatory nucleotide sequence which is employed with a selected antibody-producing cell is not critical to the method of the invention. Preferred regulatory sequences for modification of expression of immunoglobulin sequences in antibodyproducing cells include a mouse heavy chain enhancer sequence, mouse kappa (K) chain enhancer sequence, or a promoter derived from Moloney murine leukemia virus, Rous sarcoma virus or spleen focus forming virus.

Modifying sequences are inserted into recombinant DNA targeting vectors of the invention which are used to transfect antibody-producing or embryonic stem cells.

For the modification of all or a portion of a constant region of an antibody, modifying sequences of the invention may include, but are not limited to an immunoqlobulin constant region having a particular effector function, class and/or origin (e.g., IgG, IgA, IgM, IgD, or IgE constant regions of human immunoglobulins or any other species) or a portion of a constant region which modifies the activity or properties of the constant region of the immunoglobulin; as well as genes which encode other molecules that confer some new function to a modified antibody molecule, e.g., an enzyme, toxin, a biologically active peptide, growth factor, inhibitor, conjugatable peptide linker, and the like. For the modification of all or a portion of a variable region of an antibody, modifying sequences of the invention may include, but are not limited to immunoqlobulin variable regions that encode a different variable region having a different antigen affinity or specificity, or a portion of a variable region which modifies the activity or properties of the variable region of the immunoglobulin so that the resulting modified antibody has a greater affinity or higher degree of specificity for the antigen.

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The modifying sequence chosen depends, in part, upon the use intended for the modified antibody molecule expressed. For example, if therapeutic use in humans is intended, then the modifying sequence could encode a human constant region, preferably of a class having a desired effector function for the therapeutic use in mind, such as a toxin for killing targeted tumor cells. If an improvement or alteration in the existing effector function of the antibody is desired, a portion of the constant region may be replaced with a sequence that confers such improved or altered effector function to the resulting modified antibody molecule. For instance, or i

where it is desired to reduce the rate of clearance of the antibody from the circulatory systems, such as with antibodies used for imaging or delivery of cytotoxic drugs to a target cell, a portion of a human y, heavy chain region may be replaced with a region from a Y, heavy chain region containing a mutation which siminates binding of the Y, heavy chain to Fc receptors.

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of targeted delivery of an enzyme, toxin, drug, hormone or growth factor in vivo is desired, a modifying sequence encoding the enzyme, toxin, drug, hormons or growth factor or an appropriate linker for conjugation to such is used. If the modified antibodies are to be used in diagnostic assays, for example where labeled antibodies are utilized, a modifying sequence encoding an enzyme or its substrate could be used. Such enzyme/substrate systems include, but are not limited to those which produce a colored product when reacted; for example, beta-galactosidase, alkaline phosphatase, horseradish peroxidase, and the like. The resulting modified antibodies may be used as labeled antibodies in the procedures intended with or without further modification, e.g., the chemical attachment of enzymes, drugs, toxins, hormones, growth factors, and the like.

The modifying sequence used to convert antibody variable regions may comprise all or a portion of the coding sequence for a variable region of an antibody molecule that recognizes a desired antigen. These may encode antigen binding regions that recognize related or completely unrelated antigens. If an improvement or alteration in antigen binding or specificity is desired, a portion of the variable region may be replaced with a sequence that confers such improved or altered binding or specificity to the resulting modified antibody molecule. The invention method also is useful to construct cells that produce bi-specific antibodies,

i.e., antibodies which contain two different variable regions, by modifying cells to contain two different variable regions in the light and/or heavy chain genes. In this context, if a wildtype and incompatible mutant low site are integrated into the same genome, a particular modifying sequence can be inserted evaluations of these sites by using a vector which we have been as the context of these sites by using a vector when the context of these sites by using a vector when the context of these sites by using a vector when the context of th

particular modifying sequence can be inserted selectively into either of these sites by using a vector comprising a site comparible with the desired integrated site, as described, for instance, by Johnson et al., in PCT publication Now3/13172.

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cre: The enzyme expression product of the cre gene which is a recombinase that effects site-specific recombination of DNA at lox sites (see definition below). One cre gene can be isolated from bacteriophage Pl by methods known in the art, for instance, as disclosed by Abremski et al., Cell, 32:1301-1311 (1983), the entire disclosure of which is incorporated herein by reference.

Lor site: a mucleotide sequence at which the gene product of the cre gene, referred to herein as "Cre," can catalyze a site-specific recombination. A LoxF site is a 1s base pair mucleotide sequence which can be isolated from bacteriophage Fl by methods known in the art. One method for isolating a LoxF site from hatteriophage Fl is disclosed by Hoses et al., Proc. Natl. Acad. Sci. USA, 72:3398 (1982), the entire disclosure of which is hereby incorporated hartein by reference. The LoxF site consists of two 1) base pair inverted repeats separated by an 8 base pair spacer region. The mocleotide sequences of the insert repeats and the spacer region LoxF are as follows.

ATAACTTCGTATA ATGTATGC TATACGAAGTTAT

Other suitable low sites include Lox3, LoxL and LoxR
sites which are nucleotide sequences isolated from
E. coli. These sequences are disclosed and described by

Hoess et al., Proc. Natl. Acad. Sci. USA 72:1398 (1982), the entire disclosure of which is hereby incorporated herein by reference. Preferably, the lox site is LoxP or LoxC2. The mucleotide sequences of the insert remeats and the spacer region of LoxC2 are as follows.

5 repeats and the spacer region of LoxC2 are as followed academic artifacts and academic artifacts are as followed as a construction of the con

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Johnson et al., in PCT Publication W093/19172, the entire disclosure of which is hereby incorporated herein by reference, describes phage vectors in which the VM genes are flanked by two loxF sites, one of which is a mutant loxF site (loxF 511) with the G at the sewenth position in the spacer region of loxF replaced with an A, which prevents recombination within the vector from merely exciting the VM genes. However, two loxF 511 sites can recombine via Cre-mediated recombination and, therefore, can be recombined selectively in the presence of one or more widtype lox sites. The nucleotide sequences of the insert repeats and the spacer region of loxF 511 as follows.

ATAACTTCGTATA ATGTATAC TATACGAAGTTAT

Low sites can also be produced by a variety of synthetic techniques which are known in the art. For example, synthetic techniques for producing low sites are disclosed by Too et al., Nuc. Acid Mes., 12:1755 (1982) and Oglivie et al., Science, 21:42:70 (1981), the entire disclosures of which are hereby incorporated herein by reference.

Cre-Mediated Site-Specific Recombination: a process of recombination between two compatible lox sites including any of the following three events:

- deletion of a preselected DNA segment flanked by lox sites,
- inversion of the nucleotide sequence of a preselected DNA segment flanked by lox sites, and

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reciprocal exchange of DNA segments proximate to low sites located on different DNA molecules. It is to be understood that this reciprocal exchange of DNA segments results in an integration event if one or both of the DNA molecules are circular. Thus, where a loxtargeting vector is used according to the invention to insert (i.e., integrate) a modifying sequence into the genome of an antibody-producing cell having a single lox site suitable for recombination with a lox site in the vector, it is to be understood that the lox-targeting vector in such a case is comprised of a circular DNA molecule. A linear DNA segment flanked by lox sites also may be inserted into a DNA molecule comprising a single lox site by recombination between the flanking lox sites to form a circular segment followed by integration of that circle into the DNA molecule comprising a single lox site.

Targeting sequence: a sequence homologous to DNA sequences in the genome of a cell that flank or occur adjacent to the region of a immunoglobulin locus to be converted. The flanking or adjacent sequence may be within the locus itself or upstream or downstream of coding sequences in the genome of an antibody-producing cell. Targeting sequences are inserted into recombinant DNA vectors which are used to transfect antibody-producing cells such that sequences to be inserted into the cell genome, such as the sequence of a low site, are flanked by the targeting sequences of the vector.

For antibody-producing or embryonic stem cells, from the companies of the continuous continuous that direct replacement of or insertion within all or a portion of the constant region may include but are not limited to all or a portion of the V, D, J, and switch

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region (including intervening sequences called introns) and flanking sequences associated with or adjacent to the particular heavy chain constant region gene expressed by the antibody-producing cell to be transfected and may include regions located within or downstream of the constant region (including introns). Targeting sequences for light chain recombinations that direct replacement of or insertion within all or a portion of the constant region may include but are not limited to the V and J regions, their upstream flanking sequences, and intervening sequences (introns), associated with or adjacent to the light chain constant region gene expressed by the antibody-producing cell to be transfected and may include regions located within or downstream of the constant region (including introns). Targeting sequences for heavy chain recombinations that direct replacement of or insertion within all or a portion of the variable region may include but are not limited to all or a portion of the V, D, and J regions (including introns) and flanking sequances associated with or adjacent to the particular variable region gene expressed by the antibody-producing cell to be transfected. Targeting sequences for light chain recombinations that direct replacement of or insertion within all or a portion of the variable region may include but are not limited to the V and J region (including introns) and flanking sequences associated with or adjacent to the light chain variable region gene expressed by the antibody-producing cell to be transfected.

Targeting sequences for homology-targeting vectors of the present invention are usually further selected from those described above such that insertion of a lox site into a genomic sequence via site-specific homologous recombination of an associated targeting

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sequence with genomic DNA in vivo does not modify an amino acid sequence of the antibody molecule which is expressed by the transfected cell. This approach maintains the proper transcription and translation of the immunoglobulin genes which produce the desired antibody after insertion of lox sites (and, optionally, any additional sequence such as a selectable marker gene). However, in some cases it is possible to insert a lox site and other sequences into an immunoglobulin locus sequence such that an amino acid sequence of the antibody molecule is altered by the insertion, but the antibody still retains sufficient functionality for the desired purpose. For instance, Waterhouse et al., Nucleic Acids Res (1993) 21:2265-2266, the entire disclosure of which is hereby incorporated herein by reference, describes DNA sequences encoding an IgM or IgG polypeptide chain linked to a phage protein via an amino acid sequence encoded by a 34 base pair lox site. This construct produces an antibody variable region fused to the phage protein in such a way that the phage particle displays functional antibody binding sites. Functional antibody regions may be linked to other polypeptides, such as a toxin, using these or similar constructs including DNA sequences encoding an IgM or IqG polypeptide chain linked to the polypeptide via an amino acid sequence encoded by a lox site. Vector: includes plasmids and viruses and any DNA

Vector: includes plasmids and viruses and any DNA or RNA molecule, whether self-replicating or not, which can be used to transform or transfect an antibodyproducing cell.

Romology-Targeting Vector: a recombinant DNA or vector comprising a targeting sequence and any other sequence, particularly a low site and optionally a selectable marker gene, which is used to modify immunoglobuln loci using homology-mediated site

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specific recombination in antibody-producing cells or embryonic stem cells transfected with the homologytargeting vector. Homology-targeting vectors are typically transfected into antibody-producing cells in the form of linear DNA molecules to enhance the desired homologous recombination event.

Lox-Targeting Vector: a recombinant DNA or RNA

vector comprising a lox site and any other sequence, particularly a modifying sequence and optionally a selectable marker gene, which is used to modify immunoglobulin loci using Cre-mediated site-specific recombination in antibody-producing cells. The lox site of the lox-targeting vector is suitable for Cre-mediated recombination with another lox site which has been inserted into a genomic sequence of the antibodyproducing or embryonic stem cell (via a homologytargeting vector), adjacent to a region of the immunoglobulin loci which is to be modified by the modifying sequence. Integration of a modifying sequence into a single lox site in a region of the immunoglobulin loci in the genome of the antibody-producing or embryonic stem cell results in modification of that region by addition of the modifying sequence. Integration of a modifying sequence into a genome comprising two lox sites flanking a region to be modified results in replacement of that region with the modifying sequence.

Antibody-Producing Cells: The homology-targeting and low-targeting vectors are used to transfect cells that contain sequence(s) homologous to the targeting sequence of the vector and that are capable of producing immunoglobulin molecules having (a) a desired antisen specificity or affinity; (b) a desired constant region; or (c) another desired quality such as high secretion levels, large scale culture adaytability, and the like.

Such cells include, but are not limited to, cell lines derived from hybridonas which produce a sonoclonal antibody having a desirable specificity, affinity or effector function, as well as hybridona cells which themselves have sustained autations which prevent expression of heavy and/or light chains of immunoglobulin.

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In certain preferred embodiments of the invention methods, the antibody-producing cells are hybridomas derived from B-cells of mice engineered with human immunoglobulin heavy and light chain genes, as described for instance by Kucherlapati et al., in PCT Publication WO91/10741 (published July 25, 1991) and in PCT Publication WO 94/02602 (published February 3, 1994). Such mice produce human antibodies in response to immunization with any antigen normally recognized by the mouse immune system, including human antigens; and Bcells from these mice can be used to make hybridomas producing human monoclonal antibodies via conventional hybridoma production methods. In one particular embodiment using hybridomas from such transgenic mice, the genomic sequence comprising the human immunoglobulin loci in the transgenic mouse further comprises a lox site adjacent to or inserted in a region of the immunoglobulin locus which is to be modified in the hybridomas by the invention method. This lox site is inserted adjacent to or with the human immunoglobulin loci prior to their insertion into the genome of the murine embryonic stem cells by incorporation into whatever vector is used to insert the human immunoglobulin loci into the genome of the embryonic stem cells from which transgenic mice, as described, for instance, by Kucherlapati et al., supra. Conventional recombinant DNA methods are used to incorporate the lox site into this vector, adjacent to or inserted in a

region of the immunoglobulin locus which ultimately is to be modified by the invention method in an antibody producing cell obtained from the transgenic mouse.

In addition, antibody-producing cells suitable for practice of the invention methods for engineering of antibodies include host cells of both prokaryotic and sukaryotic organisms disclosed by Cabilly, et al. in U.S. Patent No. 4,816,567, the entire disclosure of which is hereby incorporated herein by reference.

The following abbreviations shall have the meanings shown below:

DHFR: dihydrofolate reductase

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qpt: guanosine phosphoryl transferase gene Neo: neomycin resistance gene

Hyg: hygromycin resistance gene

Invention methods for modifying immunoglobulin loci

The invention relates to the novel use of Cremediated site-specific recombination for engineering immunoglobulin loci, such as for class-switching of antibody genes in antibody-producing lymphoid cells in situ whereby the constant region of an antibody gene is replaced with a constant region of another class, thereby producing an antibody with a changed or "switched" isotype. The resulting cells can be used for large scale production of the modified antibodies. More particularly, the present invention provides methods to produce a cell expressing a modified antibody molecule with a modified constant region, or a lymphoid cell line expressing a modified antibody molecule, using Cremediated site-specific recombination to target gene modifications to desired locations in the genomic sequence of the immunoglobulin loci of an antibodyproducing cell.

In the methods of the invention, DNA sequences comprising one or more lox sites are first introduced into the genome of the antibody-producing cell by homologous recombination. Insertion of the lox site(s) is adjacent to a region of the immunoglobulin loci of the antibody-producing cell which is to be converted to a modified region. The cells containing a genome with one or more integrated lox sites are then transfected with a DNA molecule comprising: (a) a lox site suitable for Cre-mediated recombination with an integrated lox site, and (b) a modifying sequence. The genomic and vector lox sites are then interacted with the Cre recombinase (typically, by transient expression of the recombinase from a plasmid), thereby producing recombination between the lox sites so that the modifying sequence inserts into the genomic sequence via Cre-mediated site-specific recombination of the lox sites, thereby converting the desired portion of the genomic sequence to a modified portion.

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In this method, interacting the low sites with Cre invigically performed by transient expression of the recombinase from a plasmid. Transient interaction with Cre is particularly preferred where the object is to produce a limited number of recombination events resulting in constructs containing multiple low sites because extensive Cre-mediated recombination would tend to produce undesired deletions from such constructs. Using the method of the invention, the amino acid

sequence of an antibody molecule can be modified by transfection of an antibody-producing cell with an appropriate 'homology-targeting vector' and 'loxtargeting vector' and interaction of the low sites with the Cre recombinase. In one embodisent of the invention, a cell line that comprises immunojoulin loci emoding a desired antipen specificity or affinity

is transfected with a lox-targeting vector of the invention which comprises a recombinant DNA molecule that encodes a lox site and a "modifying sequence." The modifying sequence encodes the desired nolecule which is to replace all or a portion of the constant region of the antibody molecule expressed by the antibody-producing cell. Examples of modifying sequences for this smbodient are described above.

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vector.

In this embodiment a lox site is integrated into the genomic DNA of the antibody-producing cell using a homology-targeting vector in which the "targeting sequence" is homologous to DNA sequences found in the chromosome within or adjacent to the mature gene coding sequence for the constant region of the immunoglobulin encoded by the cell line to be transfected. After transfection, homologous recombination within the antibody-producing cell will occur; some of these recombination events will lead to the integration of the lox site into the genomic DNA of the antibody-producing cell at the desired location. Subsequent Cre-mediated site-specific recombination between a lox-targeting vector and one or more integrated lox sites results in replacement of all or a portion of the constant region of the immunoglobulin locus with the modifying sequence of the lox-targeting vector, and, therefore, the expression of modified antibody molecules by the transfected cells. The extent of replacement of the constant region gene (i.e., all or a portion) depends, of course, on the location of the integrated lox site(s) within the immunoglobulin locus of the cell and on the nature of the modifying sequence in the lox-targeting

In another aspect of the invention, a lymphoid cell line which comprises an immunoglobulin locus encoding a desired constant region is transfected with a homology-

targeting vector to insert a low site adjacent to a variable region in the genome which is to be modified, followed by transfection with a lox-targeting vector containing a modifying sequence encoding all or a portion of a variable region having a desired antigen specificity or affinity. After transfection and interacting the lox sites with Cre, Cre-mediated recombination within the lymphoid cell line will occur; some of these recombination events will lead to the replacement of all or a portion of the variable region of the immunoglobulin locus with the modifying sequence and, therefore, to the expression of modified antibody molecules by the transfered cells.

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Once the transfectant that expresses the modified antibody is identified, the practice of the invention involves culturing the transfectant and isolating the modified antibody molecules from the cell culture supernatant using techniques well known in the art for isolating monoclonal antibodies. Alternatively, the transfected cells may be cultured in sections fluid in animals and harvested using well known techniques for isolating monoclonal antibodies.

all of the modifications in immunoglobulin loci of antibody-producing cells described herein may be made by first introducing a low site into the genome of an antibody-producing cell by transfecting that antibody-producing cell by transfecting that antibody-producing cell with a homeology-targeting vector of the invention. However, antibody-producing cells with genomes comprising low sites suitable for use in the invention also may be produced by making hybridomas of B-cells from transgenic mice which contain the low site integrated into germ line DNA. These sice are produced by methods such as those described by Kucherlapati et al., in PCT Publication WOS1/10741 (published July 25, 1991) and in PCT Publication WOS1/10740 (published

PCT/US96/04445 WO 96/30498

> Pebruary 3, 1994), the entire disclosures of which are hereby incorporated herein by reference. Such transgenic mice also may be produced by other methods for production of transgenic mice which are known in the art. Generally, the embryonic stem cells used to produce the transgenic mice are transformed by any convenient method with a construct containing human immunoglobulin heavy and light chain genes and further containing a lox site adjacent to or integrated within the region of the human immunoglobulin loci (e.g., a heavy chain gene) which ultimately is to be modified in an antibody-producing cell obtained from these transgenic mics. Antibody-producing cells are obtained from the transgenic mice, for instance, by conventional hybridoma production methods.

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Use of homology-targeting vectors to insert lox sites In the methods of the invention DNA sequences

comprising lox sites are first introduced into the genome of the antibody-producing or embryonic stem cell by homology-targeted site-specific recombination using known methods for insertion of genes via homologous recombination, as described, for instance, by Fell et al. in U.S. Patent No. 5,202,238, the entire disclosure of which is hereby incorporated herein by reference. In the invention method the lox site(s) are inserted adjacent to a region of the immunoglobulin loci of the genomic sequence of the antibody-producing or embryonic stem cell which ultimately is to be converted to a 30 modified region.

The targeting vectors (i.e., homology-targeting or lox-targeting vectors) of the invention comprise recombinant DNA vectors including, but not limited to, plasmids, phages, phagemids, cosmids, viruses and the

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like, which contain the lox sites and/or modifying sequences and/or targeting sequences. As described in more detail above, the soliditying sequence may comprise any of a number of regulatory sequences or genes that encode a desired structural product whereas the targeting sequence may vary depending upon the type of antibody molecule being converted and the call-type being transfected. The targeting vectors of the invention may contain additional genes which encode selectable markers including but not limited to engymas which confer drug resistance to assist in the screening and selection of transfectants. Alternatively the vactors of the invention may be cotransfected with such markers.

Other sequences which may enhance the occurrence of recombinational events may be included in homologytargeting vectors as well. Such genes may include but are not limited to genes for either sukaryotic or procaryotic recombination enzymes such as Rec λ , topoisomerase, Rec 1 or other DNA sequences which enhance recombination such as Chi. Furthermore, sequences which enhance transcription of chimeric genes produced by homologous recombination may also be included in the vectors of the invention; such sequences include, but are not limited to, inducible elements such as the metallothionine promoter (Brinster et al., 1982, Nature 296:39-42). Various proteins, such as those encoded by the aforementioned genes may also be transfected in order to increase recombination frequencies.

For modification of antibodies, the composition of targeting sequence may vary depending upon whether the homology-targeting vector is to be used to replace all or a portion of either the variable or constant region genes of light chains or heavy chains and,

further, upon the species of the host cell to be transfected. More specifically, targeting sequences are homologous to sequences which are adjacent to or which flank the coding region for the constant or variable region, or the portion thereof, to be replaced or sitered.

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For example, in a vertebrate chromosome, mature heavy chain genes are comprised, at their 5' termini. of the VDJ regions; i.e., the variable region (V). diversity region (D), and joining region (J) followed by any remaining J regions which are not expressed (the number of J regions varies with the species), and intronic sequences. The central and 3' portion of the gene consists of the constant region exons (flanked and interspersed with intronic and untranslated sequences) which may be one of various classes (e.g., mu, delta, gamma, epsilon, alpha) each of which is associated with its own adjacent switch region. Thus, the targeting sequence used to target insertion of a lox site by homologous recombination in the heavy chain gene of an antibody-producing cell may comprise a region that is homologous to any portion of the antibody gene, depending on the desired location for insertion of the lox site. For example, the targeting sequence for inserting a lox site to direct replacement of a heavy chain constant region may comprise sequences homologous to sequences spanning any region up to and including or excluding the switch region commencing with V, D or J and would be positioned accordingly in the construction of the targeting vector; e.g., at a location 5' to the coding region of the modifying sequence. The actual targeting sequence that could be used may vary depending upon the species of the target host cell and the class of antibody expressed by the target host cell.

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In some applications the low site is inserted into the genomic sequence by homologous recombination without modifying an amino acid sequence of the antibody which ultimately is to be modified using Cre-mediated site-specific recombination. In such cases involving the mature heavy chain genes of an antibody-producing cell, a low site may be inserted into any suitably located non-translated portion of those genes, such as: at their 5' tarmini, any remaining J regions which are not expressed and intronic sequences; and in the central and 3' portion of the gene, the constant region intronic and does not disrupt expression of the desired heavy chain molecule.

By contrast to the arrangement of heavy chain antibody genes in a chromosome, the mature light chain genes are composed of a VJ region at their 5' termini, intronic sequences, and a single constant region exon. Thus, the targeting sequence used to insert a lox site by homologous recombination in the light chain gene of an antibody-producing host cell may comprise a region that is homologous to any portion of the gene, depending on the desired alteration. For example, the targeting sequence for directing the replacement of a light chain constant region may be homologous to sequences spanning all or portions of the appropriate V and J through intronic sequences preceding the coding region for the constant region of the light chain. Such targeting sequences would be appropriately positioned in the homology-targeted plasmid; e.g., at a location 5' to the lox site, such that the lox site is integrated into a non-translated sequence of the light chain genes, such as in an intronic sequences preceding the coding region for the constant region of the light chain. Once again, the actual nucleotide sequence of the targeting sequence

may vary with respect to the animal species of the target host antibody-producing cell.

In a similar fashion, targeting sequences for directing insertion of low sites for Crc-mediated replacement of heavy or light chain variable regions of antibodies may comprise sequences homologous to all or portions of the appropriate regions that flank the variable region. In any case, targeting sequences may also include coding region sequences flanking an area within an exon where only a portion of the variable or constant region is to be replaced so that the protein expressed is altered in a desired feshion.

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Hee of Cre-mediated site-specific recombination

In the methods of antibody engineering of the invention, couls containing a genome with one or more integrated lor sites are transfected with a low-targeting vector, i.e., typically a DNA molecule comprising a low site suitable for Cre-mediated site-specify recombination with an integrated low site, and the low sites are then interacted with the Cre recombinase, thereby producing recombination at the low sites. The location and orientation of the low sites determines the nature of the recombination, as described herein and more fully by Sauer et al. in U.S. Patent NO. 4,959,117, the entire disclosure of which is hereby innoorporated herein by reference.

In addition to the low site, tergeting sequence and the modifying sequence, the various targeting vectors of the invention may encode a selectable marker which assists in the screening and selection of antibodyproducing cells that have been successfully transformed. Suitable selectable marker genes include drug resistance genes, such as hyg, gpf, nee and DNFR (as defined above under "abbreviations") and the like.

Methods for introducing a DNA sequence (e.g., a low-targeting vector or a homology-targeting vector) into antibody-producing cells are known in the art. These methods typically include the use of a DNA vector to introduce the sequence into the DNA of a single or limited number of antibody-producing cells and then growing such cell or cells to generate a suitable population of cells. Preferably, the DNA sequences are introduced by a plasmid capable of transforming a selected sukaryotic cell while carrying a DNA sequence. The particular vector which is employed to introduce the DNA sequence into a selected antibody-producing cell is not critical.

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In a preferred embodiment, DNA sequences are introduced into mammalian cells according to the CaPO. transfer procedure described by Graham and van der Eb, Virology, 52:456-467 (1973), the entire disclosure of which is hereby incorporated herein by reference. Transfection of lymphoid cell lines in particular may be accomplished by any of a number of methods known to those skilled in the art, including, but not limited to calcium phosphate precipitation, electroporation, microinjection, liposome fusion, RBC ghost fusion, protoplast fusion, and the like. For homologous recombination the homology-targeting vector may be linearized by cleavage with a restriction enzyme within the targeting sequence prior to transfection in order to increase the probability of homologous recombination in the transfected cell.

In the present methods, Cre is introduced to interact with the low sites, thereby producing the sitespecific Cre-mediated recombination. In one embodiment, Cre is introduced into the cells directly by microinjection. In a preferred embodiment, the cre gene is introduced into the antibody-producing cell under the

control of a regulatory nucleotide sequence. Suitable regulatory nucleotide sequences are known in the art. The regulatory nucleotide sequence which is employed with a selected antibody-producing cell is not critical to the method of the invention. A partial list of suitable regulatory nucleotide sequences for nammalian cells includes the long terminal repeat of Molonev sarcoma virus described by Blochlinger and Diggelmann, Mol. Cell Bio., 4:2929-2931 (1984); the mouse metallothionsin-1 (MT-1) promoter described by Pavlakis and Hamer, Proc. Natl. Acad. Sci. USA, 80:397-401 (1983); the long terminal repeat of Rous sarcoma virus described by Gorman et al., Proc. Natl. Acad. Sci. USA, 79:6777-6781 (1982); and the early region promoter of SV40 described by Southern and Berg, J. Mol. Appl. Genet., 1:327-341 (1982). Sauer et al., in U.S. Patent No. 4,959,317, describes a plasmid for expression of Cre in mammalian cells, plasmid PBS31, containing a cre gene upstream from a MT-1 promoter. Activation of the MT-1 promoter with CdCl, effects expression of the cre gene in

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Since the low site is an asymmetrical nucleotide sequence, two low sites on the same DNA nolecule can have the same or opposite orientations with respect to each other. Recombinations between low sites in the same orientation result in a deletion of the DNA segment located between the resulting ends of the original DNA solecule. The deleted DNA segment forms a circular molecule of DNA. The original DNA solecule and the resulting circular molecule each contains a single low site. Recombination between low sites in opposite orientations on the same DNA molecule result in an inversion of the nucleotide sequence of the DNA segment located between the two low sites. In addition, reciprocal exchange of

cells transfected with this vector.

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DNA segments proximate to lox sites located on two different DNA molecules can occur resulting in two new recombinant DNA molecules. All of these recombination events are catalyzed by the gene product of the cre quane.

Screening and Selection of Recombinants

The ultimate test for successful targeted gene modification is the production of modified antibodies by the cell. The detection of transfectants with properly integrated vector sequences can be accomplished in a number of ways, depending upon the nature of the integrated sequences. If the targeting vector contains a selectable marker, the initial screening of transfected cells is to select those which express the marker. For example, when using a drug resistance gene, those transfectants which grow in the selection media containing the otherwise lethal drug can be identified in the initial screening. A second screening is then required to identify those transfectants which have integrated a lox site of a homology-targeting vector or a modifying sequence of a lox-targeting vector, where the latter cell will express the modified antibody.

The protocol for the second screening depends upon the nature of the inserted sequences. For example, integrated low sites are detected by treatment of isolated cell DNA with Cre in vitro followed by Southern blot analysis using appropriate restriction engress, as described, for instance, by Sauer et al., U.S. Patent No. 4,959,217. Alternatively, cells transfected with a homology-targeting vector are tested for insertion of a low site by polymerase chain reaction (PCR) amplification of the specific gene region which is supposed to contain the integrated low site and examination of the amplified DNA for the presence of a

lox site (e.g., by size or sequence), according to conventional methods. In yet another alternative approach, verification of insertion of the lox site is performed indirectly, by testing for efficiency of correct integration of a modifying sequence carried by a lox-tarcetin vector of the invention.

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The expression of a modifying sequence that encodes the constant region of a different antibody class or species is detected, for instance, by an immunoassay using antibodies specific for the particular immunoglobulin class and/or species. Alternatively, a bioassay is performed to test for a particular effector function conferred by the modifying sequence. The expression of a modifying sequence which encodes a biologically active molecule such as an enzyme, toxin, growth factor, or other peptide is assayed for the particular biological activity; for example, the transfected cell products are tested using the appropriate enzyme substrate, or target for the toxin, growth factor, hormone, and the like. Alternatively, these modifying sequence products are assayed immunologically using antibodies which are specific for the modifying sequence product.

According to a modification of a method disclosed by Fell et al. in U.S. Patent No. 5,202,218, antibody-producing cells with an integrated lox site are transfected with a lox-targeting vector comprising human constant region sequences, as described sugra, and then cultured in a single flask of salective media. After a period of the sufficient to allow selection to occur (in most cases, about 2 weeks) the surviving cells are then treated with antibody that is specific for whatever immunoglobulin constant region was originally produced by the antibody-producing cell conjugated to a toxin, such as ricin. Cells which are not successfully

transfected continue to express antibody which contains nurine constant region sequences, and are killed by the toxin conjugated to the anti-murine immunoglobulin antibody. Cells expressing modified antibodies which contain the desired human constant region sequences would survive, and are subsequently cultured in soft agar and further identified by an overlay of anti-human immunoglobulin antibody.

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The transfectants which express a modifying sequence also are tested for appropriate antigen or ligand recognition via any conventional immunological binding methods, for instance, as disclosed by Fall et al., supra or Capon et al., supra. If the cell in which antibody gens are to be modified originally is incapable of producing functional antibody (for example, because it is capable of producing only light chain), after transfection with any targeting vector of the invention, useful transfectants are identified by assays known in the art which identify functional antibody, including, but not limited to, CDC, ADCC, or inumunoprecipitation reactions.

The following examples are intended to illustrate, but not to limit, the invention.

EXAMPLE 1

"Class-Switching" of Human Antibodies Produced in Murine Hybridoma Cells

Figure 1 represents a schematic diagram of one modiment (Scheme A) of the invention method to produce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated sitespecific recombination. The cell to be modified is from a lymphoid cell line (hybridoma) with a genome containing a human constant region gene which is to be converted to a modified constant region. Such cell

lines may be generated, for instance, by transformation of murine hybridomas with vectors containing human constant region genes which are integrated using homologous recombination methods such as described by Fell, et al., supra. Such cell lines also are described by Mucherlapati et al., in PCT Publication M091/10741 (published July 25, 1991) which discloses production antigen-specific human monoclomal antibodies from mice engineered with human immunoglobulin heavy and light chain genes. B-cells from these mice are used to make hybridomas production ghuman monoclomal antibodies via conventional hybridoma production methods.

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The invention method in Scheme A requires integration of only a single lox site adjacent to the constant region gene to be modified. In this scheme, a first lox site is integrated adjacent to the constant region gene (human C,) which is to be converted to a modified constant region (human C,), via site-specific homologous recombination with genomic DNA. A vector (comprised of a circular DNA) comprising a second lox site and a selectable marker gene (DHFR) is then transfected into the cells, and the lox sites are transiently interacted with Cre. Cre is provided by cotransfection of a vector which transiently expresses a cre gene, as described above. Cre-mediated sitespecific recombination of the lox site in the genome with the lox site on the circular vector results in insertion into the genome of the modifying sequence (C,) with the selectable marker gene (DHFR). The desired transfectants, which have two lox sites flanking the DHFR and C, genes, are stable in the absence of further Cre expression and are obtained by selecting for expression of the DHFR marker gene (i.e. resistance to the drug methotrexate indicated as "Met2" in Figure 1). Verification of the desired modifications of the

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immunoglobulin loci after each transfection and selection step is provided by conventional genetic analyses, such as by PCR amplification, as described above. Figure 2 represents a schematic diagram of an

alternate embodiment (Scheme B) of the invention method to produce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated sitespecific recombination and two lox sites in the genome of the antibody-producing cell. This method comprises inserting, via site-specific homologous recombination, two lox sites flanking the constant region gene of the ganomic sequence (human C.) which is to be converted to a modified constant region (human C.). As shown in Figure 2. the cells are sequentially transfected or cotransfected with two homology-targeting vectors, one for inserting each of the two low sites. Each vector used for insertion of a lox site contains a different selectable marker gene, as illustrated in Figure 2 by use of the hyg selectable marker gene (i.e., selection for resistance to the hygromycin, "Hygh") and the neo selectable marker gene (i.e., selection for resistance to the drug G418, indicated as "G4182"). Recombinants with both lox sites inserted are obtained by selecting for both markers, either consecutively or concurrently. In any event the result of proper integration of the two homology-targeting vectors is that the two inserted lox sites flank the gene which is to be converted (C,) and

is also flanked by the inserted low sites. A vector (comprised of a circular DNA) comprising an additional low site, the modifying sequence (C,) and a third selectable marker gene (DETR) is then transfected into the cells, and the low sites are interacted vith Cre. Cre is provided by cotransfecting a vector which

each selectable marker gene used for lox site insertion

transiently expresses a cre gene, as described above. The modifying sequence inserts into the genomic sequence with the second selectable marker gene, via Cre-mediated site-specific recombination of the lox site and the vector and one or the other of the lox sites inserted in the genomic sequence. As illustrated in Figure 2, integration of the modifying sequence at the lox site adjacent to the constant region gene to be converted (Cu) is followed in some cells by excision of the first marker gene (hyg), the second marker gene (neo) and the gene to be converted (C,), via further Cre-mediated sitespecific recombination. Therefore, the desired transfectants are obtained by selecting for a transfectant expressing the third marker gene (DHFR) and not expressing the first or second marker genes, i.e., resistance to methotrexate ("Meta") and sensitivity to the drug G418 ("G418") and to hygromycin ("Hyg5), in Figure 2.

A mutated version of the dihydrofolate reductase ground in the company of the com

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EXAMPLE 2

"Class-Switching" of Human Antibodies Produced in Murine Hybridoma Cells

Comprising an Integrated Lox Site

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Figure 3 represents a schematic diagram of yet and the reproduce a cell expressing a human antibody molecule with a modified constant region using Cre-mediated site-specific recombination. The cell to be modified is from a lymphoid cell line (hybridoma) with a genome containing a human constant region gene which is to be converted to a modified constant region. The first low site is integrated adjacent to the constant region gene (human C,) which is to be converted to a modified constant region gene (numan C,) which is to be converted to a modified constant region (human C,). More particularly, Scheme C shows a lox site 5' to the C, enhancer sequences (E), the sequence of C, exor 1" (L) and C, switch sequences

shows a lox site 5' to the C, enhancer sequences (E,)
the sequence of C, exon "1" (I,) and C, switch sequences
(S,) of the human C, region.
Such cell lines may be generated, for instance, by
transformation of murine hybridomas with vectors
containing human constant region genes which are

integrated using homologous recombination methods such as described by Fell, et al., supra, where the constant region genes in the vector have been modified by insertion of the low site by any conventional genetic engineering method. Such cell lines also may be produced by making hybridomas of B-cells from transpenic mice which contain the low site integrated into gern line DNA as described hereinsbowe, by conventional hybridoma production methods.

The invention method in Scheme C requires integration of only a single low site adjacent to the constant region gene to be modified. In this scheme, a vector (comprised of a circular NDA) comprising a second low site and a selectable marker gene (e.g., DEFR) is transfected into the cells, and the lor sites are transiently interacted with Cre, as described above. Cre-mediated site-specific recombination of the low site in the genome with the low site on the circular vector results in insertion into the genome of the modifying sequence (c.) with the selectable marker gene (BUFE). The desired transfectants, which have two low sites flanking the BUFE and C, genes, are stable in the absence of further Cre expression and are obtained by selecting for expression of the DUFF marker gene (i.e. resistance to the drug methotrexate indicated as "Mat" in Figure 3). Verification of the desired modifications of the immunoglobulin loci after each transfection and selection step is provided by conventional genetic analyses, such as by PCR applification, as above.

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SEQUENCE LISTING

(1) GENERAL INFOR	MATION:
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(i) APPLICANT: JAKOBOVITS, AYA ZSEBO, KRISZTINA M.

- (ii) TITLE OF INVENTION: PRODUCTION OF ANTIBODIES
 10 USING CRE-MEDIATED SITE-SPECIFIC RECOMBINATION
 - (iii) NUMBER OF SEQUENCES: 3
 - (iv) CORRESPONDENCE ADDRESS:
- 15 (A) ADDRESSEE: CELL GENESYS, INC.
 - (B) STREET: 322 LAKESIDE DRIVE
 - (C) CITY: FOSTER CITY
 - (D) STATE: CALIFORNIA
 (E) COUNTRY: UNITED STATES
 - (F) ZIP: 94404
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 (D) SOFTWARE: PatentIn Release #1.0, Version
 - #1.25
 - (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER:(B) FILING DATE: 29-MAR-1995
 - (C) CLASSIFICATION:

-46-

SUBSTITUTE SHEET (RULE 26)

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10 (2) INFORMATION FOR SEQ ID NO:1:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (C) Discussion Daniger
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATAACTTCGT ATAATGTATG CTATACGAAG TTAT
34

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 34 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
- ACAACTICGT ATAATGTATG CTATACGAAG TTAT
 34
 - (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 34 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- 15

- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:
- 20 ATAACTICGI ATAATGIATA CTATACGAAG TTAT 34

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CLAIMS

- 1. A method to produce a cell expressing an antibody from a genemic sequence of said cell comprising a modified immunoglobulin locus, which method uses Cre-mediated 5 site-specific recombination to modify said locus and comprises.
 - transfecting an antibody-producing cell with a first homology-targeting vector comprising;
 - (i) a first lox site. and
 - (ii) a targeting sequence homologous to a first DNA sequence adjacent to the region of the immunoglobulin locus of said genomic sequence which is to be converted to a modified region, so that said first low site is inserted into said genomic sequence via site-specific homologous recombination with genomic DNA in vivo;
 - (b) transfecting said cell with a lox-targeting vector comprising:
 - (i) a second lox site suitable for Cre-mediated recombination with said first lox site, and
 - (ii) a modifying sequence to convert said region of the immunoglobulin locus to said modified region;
- 25 (c) interacting the lox sites with Cre, so that said modifying sequence inserts into said genomic sequence via Cre-mediated site-specific recombination of said lox sites, thereby converting said region of the immunoglobulin locus to said modified region; and
 - (d) selecting a transfectant in which said region of the immunoglobulin locus is converted to said modified region and which transfectant produces said antibody molecule.

 The method of claim 1 wherein said loxtargeting vactor further comprises a selectable marker gene operably linked to control regions such that said marker gene is expressed in said cell,

- so that in step (c), on interacting the lox sites with Cre, said marker gene inserts into said genomic sequence with said modifying sequence via Cre-mediated site-specific recombination of said lox sites, and
- in step (d), said selecting for a transfectant 10 comprises selecting for a transfectant expressing said marker gene.

. The method of claim 1 further comprising

an additional step before step (b) which additional step comprises transfecting said cell with a second homology-targeting vector comprising:

- (i) a third lox site suitable for Cre-mediated recombination with said first and second lox sites.
 - (ii) a first selectable marker gene operably linked to control regions such that said first marker gene is expressed in said cell, and

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- (iii) a targeting sequence hosologous to a second DNA sequence adjacent to the region of the immunoglobulin locus of anid genomic sequence which is to be converted, anid region which is to be converted being flanked by said first and second DNA sequences, so that eaid third low site and said first marker gene are inserted into said genomic sequence vis site-specific hosologous recombination with genomic DNA in vivo; and
- 20 wherein in step (b), said lox-targeting vector further comprises a second selectable marker gene operably linked to control regions such that said second marker gene is expressed in said cell,
- so that in step (c), on interacting the lox sites with 25 Cre, said second marker gene inserts into said genomic sequence with said modifying sequence via Cre-mediated site-specific recombination of said lox sites; and
- in step (d), said selecting for a transfectant comprises selecting for a transfectant expressing said 30 second marker gene.

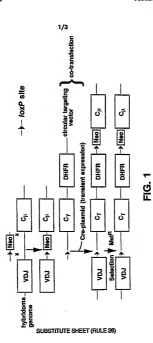
The method of claim 3 wherein

in step (c), on interacting the low sites with Cre, after said modifying sequence and second marker gene insert into said genomic sequence, said first marker gene and said 5 region to be convexted are deleted by Cre-mediated site-specific recombination; and

in step (d) said selecting for a transfectant further comprises selecting for a transfectant not expressing said first marker gene.

- 5. The method of claim 1 wherein said region of the immunoflobulin locus to be converted to a modified region comprises a constant region gene and said modifying sequence comprises one of:
- a) a regulatory nucleotide sequence which replaces 15 all or a portion of the regulatory sequences of the immunoglobulin locus of said genomic sequence to provide modified expression of said immunoglobulin locus;
- b) a nuclectide sequence that encodes a translation product to replace all or a portion of either the constant region or the variable region of an antibody molecule to form a modified antibody molecule; and
 - a sequence encoding an enzyme, toxin, hormone, growth factor or linker.
- The method of claim 5 in which said constant
 region gene of said genomic sequence comprises one of:
 - a murine constant region gene and said modifying sequence comprises a human constant region gene or portion thereof;
- b) a human constant region gene and said modifying 30 sequence comprises a different human constant region gene or portion thereof; and
 - c) a human Cy gene and said modifying sequence comprises a human Cy constant region gene.

- The method of claim 1 wherein said cell expressing an antibody molecule is a murine hybridoms cell line producing a human antibody by expression of a human immunoclobulin locus.
- 5 8. A method to produce an antibody from a cell expressing said antibody from a genomic sequence comprising a modified immunoglobulin locus wherein said cell is obtainable by the method of claim 1, which method comprises:
 - (a) incubating said cell under conditions such that said cell expresses said antibody, and
 - (b) recovering said antibody expressed by said cell.
- An embryonic stem cell of a non-primate mammal
 comprising a genome comprising:
 - at least a functional portion of a human heavy chain immunoglobulin locus or at least a functional portion of a human light chain immunoglobulin locus,
- wherein a lox site is adjacent to or integrated within 20 a region of said human heavy chain or light chain immunoglobulin locus.
 - 10. A transgenic non-primate mammal comprising a genome comprising:
- at least a functional portion of a human heavy chain 25 immunoglobulin locus or at least a functional portion of a human light chain immunoglobulin locus,
 - wherein a lox site is adjacent to or integrated within a region of said human heavy chain or light chain immunoglobulin locus.



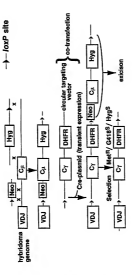


FIG. 2

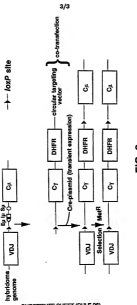


FIG. 3

INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04445

	CLASSIFICATION	OF SUBJECT	MATTER

IPC(6) :C12N 5/00, 5/06, 5/16, 15/00, 15/06, 15/09, 15/13

US CL :435/172.3, 240.21; 800/2

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/172.3, 240.21; 800/2

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

Databases: STN, Biosis, CA Search Terms: Jakobovits/Jau; zsebo7/au; homolog?; recomb?; cre; lox; es; embryonic; stem; immunoglob?; heavy: light; antibod?; immuns: constant; variab?

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Υ	GU, H. et al., Independent control of immunoglobulin switch recombination at individual switch regions evidenced through Cre-LoxP-mediated Gene Targeting, Cell. 18 June 1993, Vol. 73, pages 1155-1164, see entire document.	1-10
Y	CAPECCHI, M.R., Altering the genome by homologous recombination. Science. 16 June 1989, Vol. 244, pages 1288-1292, see entire document.	1-10
Y	WO 94/02602 A1 (CELL GENESYS, INC.) 03 February 1994, see entire document.	1-10
Υ	US 5,202,238 A (H.P. FELL, JR. ET AL.) 13 April 1993, see entire document.	1-10

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A*	document defining the general state of the set which is not considered to be of particular colorance.		practice of theory underlying the investion
		X	document at pretable relevance; the chiral inventors cannot be
2.	carior document published on or other the unconstrood filling date		considered moved or connect be considered to provide no inventive step
1.5	Jacobson which may show dealth on priority choosing or which w		when the document is taken where
-	used to extellish the publication date of souther citation or other special reason (as specified)	-Y-	decument of purcuedor extensors; the claimed severation cannot be completed to accorde an associate step when the document is
υ.	decrease referring to an unit electronic, use, exhibition or other more	comband with one or more utiled as the deceatests. So being obvious to a person skilled as the art	continued with one or more until much discentifie, such combination being obscur to a person skilled as the act
T*	document published prior to the international filling duce but beer than	.z.	document member of the name point family

Date of the actual completion of the international search

15 JULY 1996

Date of mailing of the international search

29 AUG 1996

Further documents are finted in the continuation of Box C. See potent family annex.

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/04445

Category*	mustion). DOCUMENTS CONSIDERED TO BE RELEVANT Lating of document, with indication, where appropriate, of the relevant passages Relevant to claim No.				
Category*	Cuation of document, with stockaston, where appropriate, of the resevant pussages	Resevant to cumin 140			
Y	US 4,959,317 A (B.L. SAUER) 25 September 1990, see entire document.	1-10			
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	-				